A Review on Genetic Dominant Disorder-Polydactyly

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ABSTRACT
Polydactyly is genetic disorder in which there is mutation of gene that is located on short arm of chromosome 7. One gene that can cause polydactyly is GLI3 and it is one among number of genes that are known to be involved in the patterning of tissues & organs during development of the embryo. Mutations of GLI3 gene during development will cause two types of polydactyly. Those are postaxial(ulnar) and preaxial(radial) polydactyly.

The treatment plan is based on the outcome of analysis of patient's medical history, social history, motivation, social and psychological disturbance.

KEYWORDS: Mutations, gene, postaxial, preaxial, polydactyly

INTRODUCTION
A genetic disorder is a genetic problem caused by one or more abnormalities in genome, especially a condition that is present from birth. Genetic disorders may be hereditary, passed down from the parents' genes. In genetic disorders, defects may be caused by new mutations or changes in DNA sequence. In such cases, the defect will only be passed down if it occurs in the germ line. Polydactyly is one such genetic disorder in which, the mutation of gene that is located on short arm of chromosome 7 is seen. One gene that can cause polydactyly is GLI3 and it is one among number of genes that are known to be involved in the patterning of tissues and organs during development of the embryo. It does this by helping to control whether specific genes are turned on or off. That is because GLI3 is a transcriptional repressor, which means that it codes for protein (in fact a DNA-binding protein) that regulates the expression of one or several genes by decreasing the rate of transcription. Mutations of GLI3 gene during development are known to cause two types of polydactyly. Those are preaxial and postaxial polydactyly.

Development of limb usually takes place proximodistally, once positioning along the craniocaudal axis is determined, growth must be regulated along the proximodistal, anterioposterior, and dorsoventral axis. It is initiated by TBX5 and FGF10 in the forelimb and TBX4 and FGF10 in the hindlimb secreted by lateral plate mesodermal cells.

CASE REPORT
A foetal specimen of polydactyly present in the museum of department of Rachana Shareera, Sri Dhamasthala Manjunatheshwara College of Ayurveda & Hospital, Hassan was selected for observational study.

Specimen: A male foetus showed postaxial or ulnar polydactyly in both hands
There are three types of classification of polydactyly they are preaxial or radial, postaxial or ulnar and central. Among them preaxial is most common, refers to duplication of first digital rays.4

The foetal specimen found in museum of department of rachana shareer, SDMCA & H, Hassan it was found that the male foetus had postaxial polydactyly of both upper limbs.

DISCUSSION

Polydactyly follows an autosomal dominant pattern of inheritance which means that

- It is not sex linked, so that boys and girls may be equally affected and
- A child who has polydactyly parent has 50% chance of being polydactyly.

Polydactyly is reported to occur in 1 in every 500 livebirths and hence it is not uncommon.

Positioning of the limbs along the craniocaudal axis in the flank region of the embryo is regulated by the HOX gene expressed along this axis. These homeobox genes are expressed in overlapping patterns from head to tail with some having more cranial limits than others. For example, the cranial limit of expression of HOXB8 is at the cranial border of forelimbs, and misexpression of this gene alters the position of limbs.

Once outgrowth is initiated, bone morphogenetic proteins (BMPs), expressed in ventral ectoderm, induce formation of apical endodermal ridge (AER) by signalling through the homeobox gene MSX2. Expression of radical fringe, in the dorsal half of the limb ectoderm, restricts the location of AER to the distal tip of the limbs. This gene induces expression of SER2, a homologue of drosophila serrate, at the border between cells that are expressing radical fringe and those that are not. It is at this border that the AER is established. Formation of the border itself is assisted by expression of engrailed-1 in ventral ectoderm cells, because this gene represses expression of radical fringe. After the ridge is established, it expresses FGF4 and FGF8, which maintain the progress zone, the rapidly proliferating population of mesenchyme cells adjacent to the ridge. Distal growth of limb is then affected by this rapidly proliferating cells under the influence of the FGF’s. As growth occurs, mesenchymal cells at the proximal end of the progress zone become farther away from the ridge and its influence begin to slow down their division rates and differentiation.

Patterning of the anteroposterior axis of the limb is regulated by the zone of polarizing activity (ZPA), a cluster of cells at the posterior border of the limb near the flank. These cells produce retinoic acid (vitamin A), which initiates expression of sonic hedgehog (SHH), a secreted factor that regulates the anteroposterior axis.

Thus, for example, digits appear in the proper order, with the thumb on the radial (anterior) side. As the limb grows, the ZPA moves distal ward to remain in proximity to the posterior border of the AER. Misexpression of retinoic acid or SHH in the anterior margin of the limb containing a normally expressing ZPA in the posterior border results in mirror image duplication of limb structures.
The dorsoventral axis is regulated by BMP’s in the ventral ectoderm. Which induce expression of the transcription factor EN1. In turn, EN1 represses WNT7a expression, restricting it to the dorsal limb ectoderm. WNT7a is a secreted factor that induces expression of LMX1, a transcription factor containing a homeodomain, in the dorsal mesenchyme. LMX1 specifies cells to be dorsal, establishing the dorsoventral components. In addition, WNT7a maintains SHH expression in the ZPA and therefore indirectly affects anteroposterior patterning as well. These two genes are also intimately linked in signalling pathways in Drosophila, and this interaction is conserved in vertebrates. In fact, all of the patterning genes in the limb have feedback loops. Thus, FGF’s in the AER activate SHH in the ZPA, while WNT7a maintains the SHH signal.

Although patterning genes for the limbs axes have been determined, it is the HOX genes that regulate the types and shapes of the bones of the limb. Thus, HOX gene expression, which results from the combinatorial expression of SHH, FGFs, and WNT7a, occurs in phases in three places in the limb that correspond to formation of the proximal (stylpod), middle (zeugpod), and distal (autopod) parts. Genes of the HOXA and HOXD clusters are the primary determinants in the limb, and variations in their combinatorial patterns of expression may account for differences in forelimb and hindlimb structures. Just as in the craniocaudal axis of the embryo, HOX genes are nested in overlapping patterns of expression that somehow regulate patterning. Factors determining forelimb versus hindlimb are the transcription factors TBX5 (forelimb) and TBX4 together with PITX1 (hindlimbs)⁶.

CONCLUSION
Polydactyly is the most common congenital anomaly of the hand and foot. The frequency of polydactyly varies widely among populations. It may be an isolated condition or part of a congenital syndrome. Polydactyly is generally classified into three major groups: medial ray (preaxial), central ray and lateral ray (postaxial).

The deformity is cosmetically unacceptable in many cultures and this in turn leads to psychological burden. Polydactyly often causes social embarrassment and results in compromised to move the hand independently. It often has severe impact on patients self-esteem and hampers his or her quality of life. Hence the treatment plan is formulated and tailored specifically for each individual patient. Though evaluation and diagnosis are among most important aspects of overall patient's management, the treatment plan is based on the outcome of analysis of patient's medical history, social history, motivation, and psychological disturbances. A patients ultimate satisfaction with treatment outcome often depends upon attention to the patients’ chief concerns⁷. In the dead foetal specimen mounted in museum of SDMCA & H, Hassan, it was observed that the polydactyly was of post axial or ulnar polydactyly in both upper limbs.

Radiographs of the affected limb are recommended to show whether the rudimentary digit contains skeletal elements. The degree of deviation of the digit and the size of the articulating metacarpal or metatarsal may also be helpful and surgical planning.

REFERENCES
[4] Left handed polydactyly: a case report, Nicola mumali et all